



Veeva Announces eSource Application for Research Sites to Eliminate Paper and Streamline Clinical Trial Data Flow

Descrizione

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Frees sites from duplicate data entry and connects EHR and EDC for higher quality trial data, faster

PLEASANTON, Calif., Jan. 29, 2026 /PRNewswire/ - Veeva Systems (NYSE: VEEV) today announced Veeva eSource, a new Veeva SiteVault application designed to significantly reduce manual clinical trial processes and increase data quality. Veeva eSource will eliminate paper at the site through a direct data capture application and streamline data through EDC integration, and EHR to EDC transfer.

"I'm excited to see Veeva eSource in action; although there are other options in use by sites, traditional eSource is still incredibly time-consuming to program, complete, and transfer data from eSource to EDC," said Alisha Garibaldi, CEO, Skylight Health Research. "A seamless flow of data from Veeva eSource to EDC will reduce errors, minimize QC processing, and allow us to spend more time where it matters - with our patients."

With Veeva eSource, clinical trial data flows from site to sponsor with less effort through:

"For the first time, we are connecting data and processes to enable straight-through clinical data flow from site to sponsor," said Jim Reilly, president of Veeva Development Cloud. "It is a major step toward our vision to simplify, standardize, and connect clinical trials for higher efficiency and a better experience for sponsors, sites, and patients."

"Veeva eSource advances our goal of delivering significant efficiency and simplicity for clinical trial sites," said Nick Frenzer, general manager of Veeva site solutions. "By delivering eSource, we can complete the picture to provide integrated data transparency across a site to improve trials."

Veeva eSource is part of the Veeva SiteVault platform and requires SiteVault CTMS at the site to simplify and streamline the visit experience for the site and patient. Veeva eSource is planned for early adopter availability in the second half of 2026.

About Veeva Systems Veeva delivers the industry cloud for life sciences with software, data, and business consulting. Committed to innovation, product excellence, and customer success, Veeva serves more than 1,500 customers, ranging from the world's largest biopharmaceutical companies to emerging biotechs. As a Public Benefit Corporation, Veeva is committed to balancing the interests of all stakeholders, including customers, employees, shareholders, and the industries it serves. For more information, visit veeva.com.

Veeva Forward-Looking Statements This release contains forward-looking statements regarding Veeva's products and services and the expected results or benefits from use of our products and services. These statements are based on our current expectations. Actual results could differ materially from those provided in this release and we have no obligation to update such statements. There are numerous risks that have the potential to negatively impact our results, including the risks and uncertainties disclosed in our filing on Form 10-Q for the period ended October 31, 2025, which you can find here (a summary of risks which may impact our business can be found on pages 33 and 34), and in our subsequent SEC filings, which you can access at sec.gov.

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